

ATGAGAAGGTGTAGAATAAGTGGGAGGCCCCCGGCGCCCCCGGTGTCCCCGCCCAGGCC MetArgArgCysArgIleSerGlyArgProProAlaProProGlyValProAlaGlnAla

 ${\tt CCTGTCTCCCAGCCTGATGCCCCTGGCCACCAGAGGAAAGTGGTGTCATGGATAGATGTGProValSerGlnProAspAlaProGlyHisGlnArgLysValValSerTrpIleAspVal}$

TATACTCGCGCTACCTGCCAGCCCCGGGAGGTGGTGGTGCCCTTGACTGTGGAGCTCATG TyrThrArgAlaThrCysGlnProArgGluValValValProLeuThrValGluLeuMet

 $\label{thm:control} TGCCCTGACGATGGCCTGGAGTGTGCCCACTGGGCAGCACCAAGTCCGGATGCAGATC \\ CysProAspAspGlyLeuGluCysValProThrGlyGlnHisGlnValArgMetGlnIle \\$

TGTGAATGCAGACCTAAAAAAAAGGACAGTGCTGTGAAGCCAGACAGGGCTGCTACTCCC CysGluCysArgProLysLysAspSerAlaValLysProAspArgAlaAlaThrPro

CACCACCGTCCCCAGCCCCGTTCTGTTCCGGGCTGGGACTCTGCCCCCGGAGCACCCTCC HisHisArgProGlnProArgSerValProGlyTrpAspSerAlaProGlyAlaProSer

 ${\tt CCAGCTGACATCACCCAATCCCACTCCAGCCCCAGGCCCCTCTGCCCACGCTGCACCCAGProAlaAspIleThrGlnSerHisSerSerProArgProLeuCysProArgCysThrGln}$

 ${\tt CACCACCAGTGCCCTGACCCCGGACCTGCCGCTGCCGCTGTCGACGCCGCAGCTTCCTC}\\ His {\tt HisGlnCysProAspProArgThrCysArgCysArgCysArgArgArgArgSerPheLeu}\\$

CGTTGTCAAGGGCGGGCTTAGAGCTCAACCCAGACACCTGCAGGTGCCGGAAGCTGCGA ArgCysGlnGlyArgGlyLeuGluLeuAsnProAspThrCysArgCysArgLysLeuArg

AGGTGA ArgEnd

FIGURE 1 1/1

10 20 30 40 MNFLLSWVHWSLALLLYLHHAKWSQAAPMAEGGGQNHH-EVVKFMDVYQRSYC ...||...|. .||...||| |-. MRRCRISGRPPAPPGVPAQAPVSQPDAPGHQRKVVSWIDVYTRATC 60 70 80 90 100 HPIETLVDIFQEYPDEIEYIFKPSCVPLMRCGGCCNDEGLECVPTEESNITMOIMRIKPH QPREVVVPLTVELMGTVAKQLVPSCVTVQRCGGCCPDDGLECVPTGQHQVRMQILMIR-Y 120 130 140 150 QGQHIGEMSFLQHNKCECRPKK-----DRA----RQEKKSVRGKGKGQKRKRKKSRY PSSQLGEMSLEEHSQCECRPKKKDSAVKPDRAATPHHRPQPRSVPGWDSAPGAPSPADIT 190 KSWSVPCGPCSERRKHLFVQDPQTCKCSCKNTD-SRCKARQLELNERTCRCDKPRR QSHSSPRPLCPRCTQHHQCPDPRTCRCRCRRSFLRCQGRGLELNPDTCRCRKLRR

Figure 2. Sequence alignment of VEGF3 (lower line) compared to VEGF from human (upper line).

FIGURE 2 1/1